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Docket No. 47,653.1 (1789)

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: J.C. Houck
Serial Number: 09/189,130 Art Unit: 1654
Filed: November 10, 1998 Examiner: M. Borin
For: SMALL PEPTIDES AND METHODS FOR TREATMENT OF
ASTHMA AND INFLAMMATION

#13
Plunkett
4/19/00

CERTIFICATE OF EXPRESS MAILING

I hereby certify that this paper (along with any paper referred to as being attached or enclosed) is being deposited with the United States Postal Service on April 5, 2000, 2000, in an envelope addressed to the Assistant Commissioner for Patents, Washington, D.C. 20231.

By: Barbara J. James

Honorable Assistant Commissioner for Patents
Washington, DC 20231

Sir:

DECLARATION OF JAMES CLAGETT

I, James Clagett, hereby declare that:

1. I am a citizen of the United States of America residing 5615 139th Avenue SE, Snohomish, Washington 98290.

2. I hold a Ph.D. in microbiology from the University of Nebraska. I have over 30 years experience in research and development related to microbiology, particularly in the fields of immunology and immunopathology. A copy of my curriculum vitae is attached hereto as Attachment A.

3. Since 1997, I have been a consultant providing scientific expertise to

the biotechnology and pharmaceutical communities.

4. I personally performed or directly oversaw the experiments which produced the results presented and discussed herein.

5. I have read and understand the Office Action of January 12, 2000, including the references cited therein.

6. The present invention is directed to a pharmaceutical composition having anti-inflammatory activity comprising novel formyl methionyl peptides having the formula f-Met-Leu-X, wherein X is selected from the group consisting of Tyr, Tyr-Phe, Phe-Phe and Phe-Tyr.

7. Based on my knowledge and experience in the field, it is my opinion that it is well known to those skilled in the art that formyl methionyl peptides have pro-inflammatory activity. Specifically, it is well documented in the literature that the fMLP, and other prior art peptides, stimulate conditions associated with inflammation, namely mast cell degranulation and chemotaxis to the site of inflammation.

8. I have performed the following experiments.

Briefly, female Balb/CJ mice were injected subcutaneously into the dorsum of the feet with either (a) 10 μ g fMLP peptide, (b) 200 μ g of fMLP, (c) 200 μ g fMLP and 200 μ g FMLPP simultaneously or (d) vehicle alone (4% v/v ? DMSO in Tyrode's solution) as a control. The animals were sacrificed at 10 or 30 minutes post-injection and the feet collected for histological examination. The cutaneous soft tissues were dissected from the feet and embedded in paraffin. 5-7 μ sections were cut and stained with H&E (what is this?) for the detection of cellular content and location within the muscularis and dermis.

The results are illustrated in figures in Attachment B.

9. Figure 1 shows stained tissue sections harvested from mice 10 minutes after injection with 10 μ g of fMLP into the subcutaneous layers of the skin on the dorsum of the feet. Polymorphonuclear cells (PMNs) have migrated into the interstitium and have attached to vessel walls.

Panel A (x160) shows numerous PMNs (arrows) located within the intercellular matrix and the connective tissue appears edematous and slightly hemorrhagic.

Panel B (x160) shows the extravascular migration of leukocytes from an oblique section of a small blood vessel in the muscularis region.

Panel C (x320) shows the extravascular migration of neutrophils from another blood vessel.

10. Figure 2 compares stained tissue sections harvested from mice 30 minutes after injection with (i) 200 μ g of fMLP alone, (ii) 200 μ g fMLP and 200 μ g fMLPP simultaneously or (iii) vehicle alone into the subcutaneous layers of the skin on the dorsum of the feet.

Panel A (x100) shows the results of injection with 200 μ g of fMLP alone. A massive cellular infiltration and a reddish reaction material is observed in the interstitium of the skin. Many leukocytes are in the extravascular spaces and associated with small blood vessels (arrows).

Panel B (x100) shows the results of simultaneous injection with 200 μ g of fMLP and 200 μ g of fMLPP. fMLPP reduced the cellular infiltration observed when fMLP was injected alone (see panel A). The vessels have no PMNs inside or outside (arrows).

Panel C (x100) shows the results of tissues harvested from control animals injected with vehicle (4% DMSO) alone. No cellular infiltration is seen and no PMNs are observed in association with the small blood vessels (arrows).

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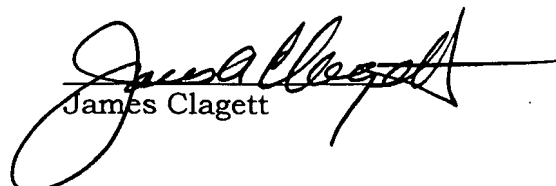
11. From these experiments, it can be concluded that fMLP (prior art) administration into the skin of mice produced an intense vascular and extravascular accumulation of inflammatory cells, largely neutrophils. Simultaneous administration of fMLPP (invention) with fMLP (pro-inflammatory) reduced the inflammatory response to near normal levels.

12. Based on my knowledge and experience in the field, it is my opinion that one skilled in the art would readily conclude from these experiments that the prior art peptide, fMLP, has pro-inflammatory activity and that the peptide of the current invention, fMLPP, has anti-inflammatory activity.

13. Based on my knowledge and experience in the field, it is my opinion that the surprising and unexpected anti-inflammatory results achieved with the peptides of the present invention would not have been obvious to those skilled in the art.

14. I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and, further, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Codes, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

3/29/2000
Date


James Clagett

JAMES A. CLAGETT, Ph.D.
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EXPERIENCE:

CLAGETT CONSULTING
5615 139TH AVE SE
SNOHOMISH, WASHINGTON, 98290

Founder and Partner: January 1, 1997 to present.

Providing quality expertise to the biotechnology and pharmaceutical communities. Lead an assembled group of individuals with experience in regulatory affairs, preclinical and clinical studies. The team has worked together for an average of 3 years and was instrumental in bringing products from research through production for GenSci Regeneration Laboratories Inc., Irvine, CA. Current list of clients includes Histatek, LLC, San Francisco, CA and BioTherapeutic Computers, Seattle, WA.

BIOCOLL LABORATORIES INC.
562 1ST AVENUE SOUTH
SEATTLE, WASHINGTON, 98104

Vice President and Scientific Director: January 1993 to August 1, 1997.

Responsible for implementing and maintaining the company's overall scientific plan. Initiated the development of the company's first product, the Tissue Bone Matrix sponge and the DynaGraft family of products and human banked tissue. Supervised the writing and implementation of the Standard Operating procedures for the TBM sponge. Initiated the scientific research of new products and managed numerous consultants supporting the company's scientific strategic plan. Carried the company from conceptual design of products through introduction to the marketplace including continuous interaction with FDA.

DENTAL DIAGNOSTIC SERVICES
2000 116TH AVENUE NE.
BELLEVUE, WASHINGTON 98004

Founder and Officer: December 1992 through January 1, 1993

Responsible for taking the company from no revenues to profitability in one year. Company had over 450 clients using the sterility testing services of Dental Diagnostic Services.

THE DENTAL RESOURCE INC.
2000 116TH AVENUE NE,
BELLEVUE, WASHINGTON 98004

Consultant: October 1990 to December 1992

Responsible for developing product concept for novel water filtration device for dental offices. Participated heavily in selling devices to dental practitioners.

ULTRA DIAGNOSTICS CORPORATION
4526 11th Avenue NE, Seattle, Washington, 98108

Chief Executive Officer, President, and Scientific Director: December 1988 to December 1989.

Responsible for implementing and maintaining the company's overall business plans. Maintained on-going communications with investors. Supervised the recruitment of research and development personnel and management finance, administration, marketing and regulatory functions. Responsible of business development where a contract of \$500,000 for research and development of new reagents was concluded. Responsible for continued venture funding of approximately \$900,000. Maintained ongoing contact with key administrative and upper management individuals of the biotechnology and academic sectors in the Northwest.

President and Scientific Director: December 1984 to December 1988. Implemented the company's research and development plans and assisted the Chief Executive Officer with business development functions where research and development contracts worth \$500,000 were secured. Coordinated the use of the Scientific Advisory Board and recruited and hired key scientific personnel. Assisted the Chief Executive Officer in capital formation from "seed" through venture financing. The capital raised was in excess of \$4 million.

**UNIVERSITY OF WASHINGTON
Schools of Medicine and Dentistry
Seattle, Washington, 98105**

Affiliate Professor: 1987 to 1989.

Taught a course in Immunology and Immunopathology to Graduate Dental Students.

Professor of Periodontics and Microbiology and Immunology: 1983 to 1987

Associate Professor of Periodontics and Microbiology and Immunology: 1978 to 1983.

Research Assistant Professor of Periodontics and Microbiology and Immunology: 1973 to 1978

Taught graduate and undergraduate Immunology and Immunopathology to Microbiology and Immunology majors as well as participated in team teaching to dental and medical students. Received well above average reviews of teaching skills by students. Maintained a research laboratory with technicians, 2 doctoral and 6 masters' students. Obtained research support averaging \$150,000 per year and served on the Faculty Senate and the Human Subjects Review Committee. Served as a member on 15 doctoral and masters thesis committees.

PROFESSIONAL ASSOCIATIONS:

Editorial Board of Journal of Dental Research
Biomedical Research Support Grant Committee
American Association of immunologists
Sigma Xi

EDUCATION:

Scripps Clinic and Research Foundation	Postdoctoral Fellowship	1973
University of Nebraska	Ph.D.	1970
DePauw University	BS	1964

PUBLICATIONS

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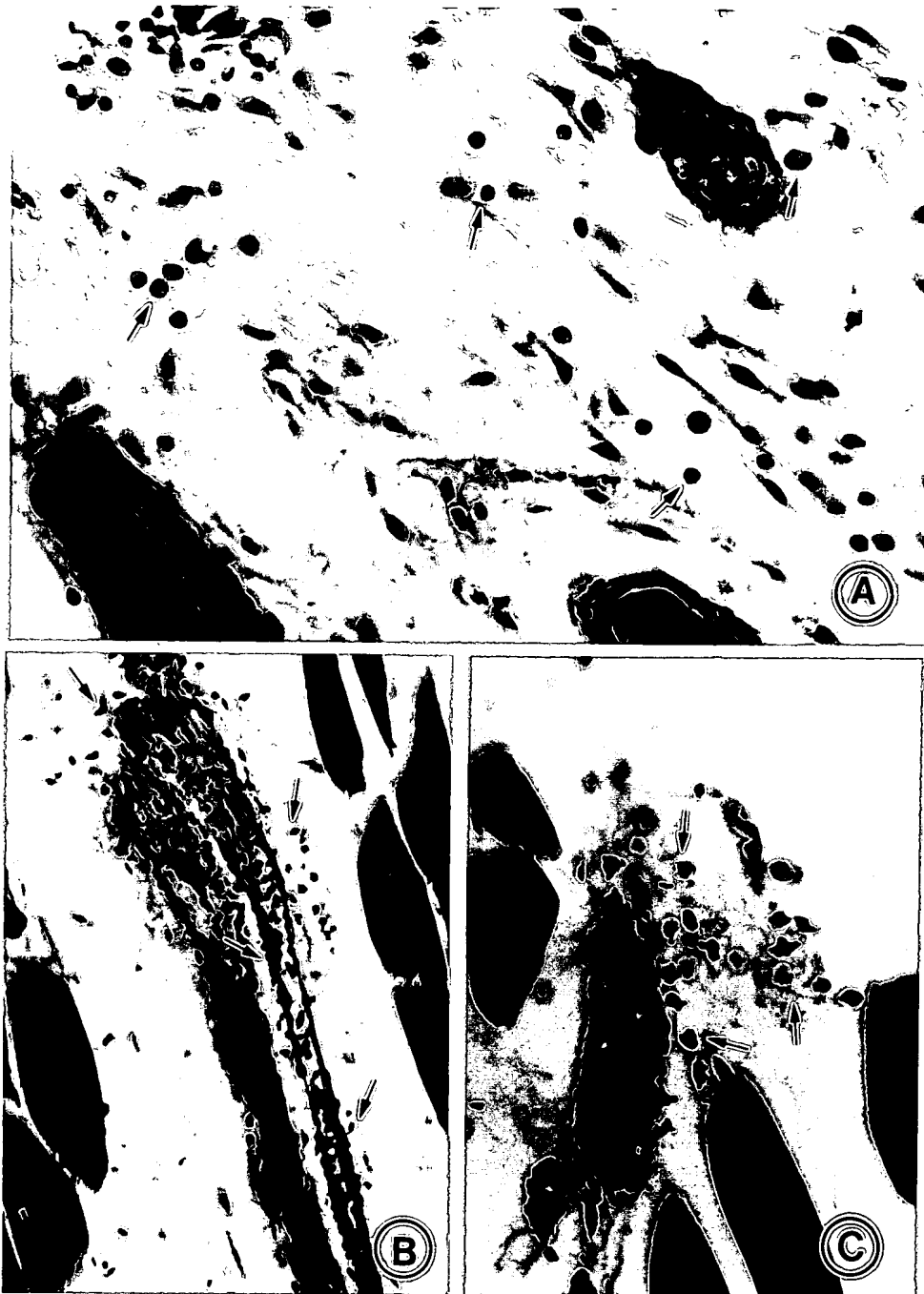


FIG. 1



FIGURE 1

Ten minutes after injection of 10 ug of fMLP into the subcutaneous layers of the skin on the dorsum of mice feet, PMNs have migrated into the interstitium and have attached to vessel walls.

PANEL A. Numerous PMNs (arrows) are located within the intercellular matrix and the connective tissue appears edematous and slightly hemorrhagic. X160

PANEL B. An oblique section of a small blood vessel in the muscularis region shows the extravascular migration of leukocytes (arrows). X160

PANEL C. At higher magnification, another vessel shows the diapedesis of neutrophils (arrows). X320

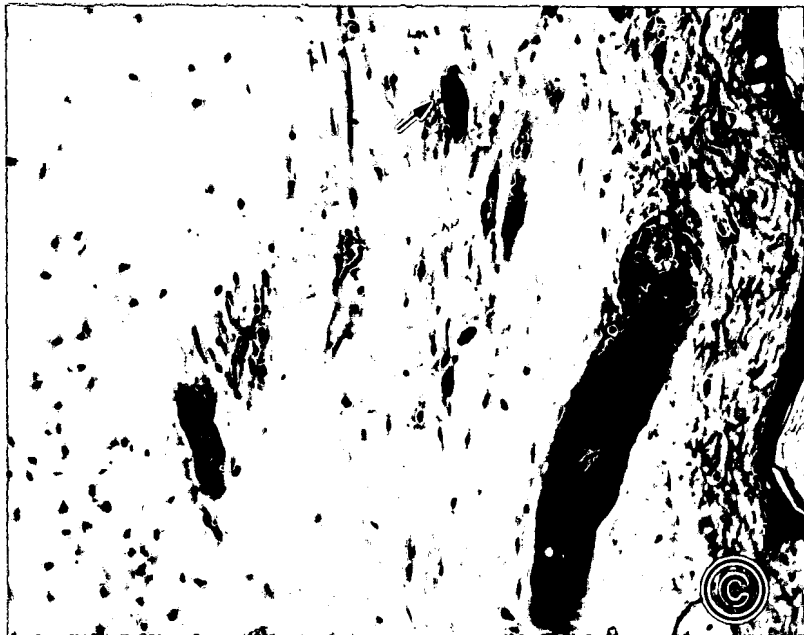
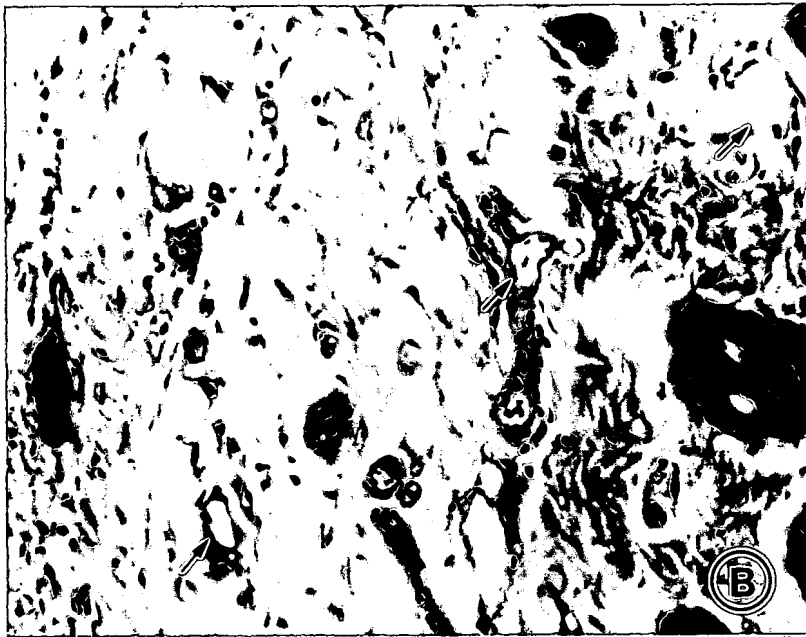
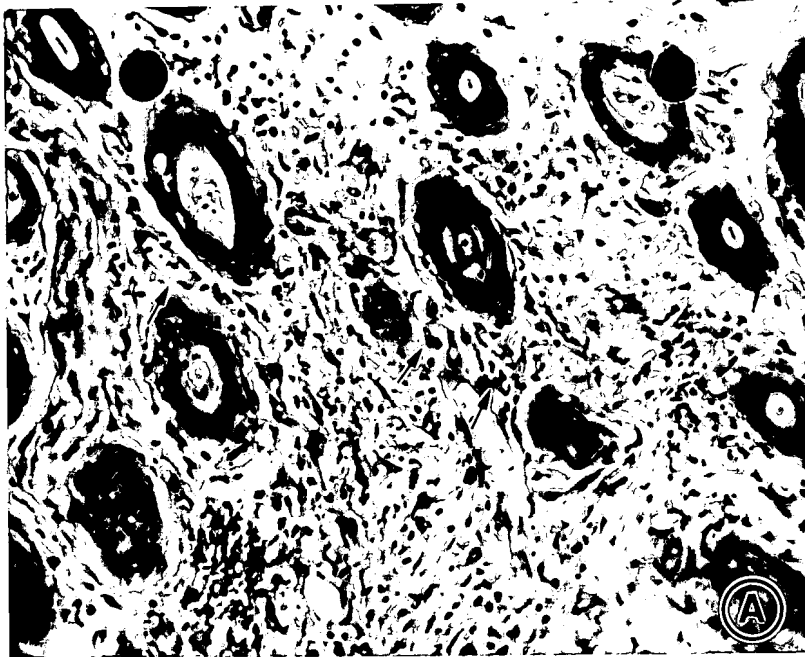


FIG. 2



FIGURE 2

Comparison of the induction of PMN diapedesis in the dermal tissues of mice when injected with HK-X simultaneously with fMLP.

PANEL A. Thirty min after injection of 200 ug of fMLP, a massive cellular infiltration and a reddish reaction material is observed in the interstitium of the skin. Many leukocytes are in the extravascular spaces and associated with small blood vessels (arrows). X100

PANEL B. The simultaneous injection of HK-X reduced the cellular infiltration observed when fMLP was injected alone (See panel A). The vessels have no PMNs inside or outside (arrows). X100

PANEL C. Tissues harvested from an animal which received only vehicle show no cellular infiltration. No PMNs are observed in association with the small blood vessels (arrows). X100